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- (71) Applicant (for all designated States except US): TEX-CONTOR ETABLISSEMENT [LI/LI]; Heiligkreuz 40, Postfach 39, FL-9490 Vaduz (LI).
- (72) Inventor; and
- (75) Inventor/Applicant (for US only): CID, Pau [ES/ES]; Rolabo SL, Calle Balmes, 85,3, E-08080 Barcelona (ES).
- (74) Agent: FRANK B. DEHN & CO.; 179 Queen Victoria Street, London EC4V 4EL (GB).

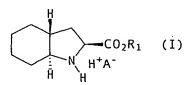
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(54) Title: A METHOD FOR THE PREPARATION OF (2S, 3AR, 7AS)-OCTAHYDRO-1H-INDOLE-2-CARBOXYLIC ACID AS KEY INTERMEDIATE IN THE PREPARATION OF TRANDOLAPRIL BY REACTING A CYCLOHEXYL AZIRIDINE WITH A DIALKYL MALONATE



(57) Abstract: A method for the synthesis of a compound of formula (I) as a mixture of enantiomers, formula (I) (wherein R_1 is H or an acid protective group and H+A indicates an optional acid with which the compound of formula (I) may form an ammonium salt) said method comprising; A) reacting a cyclohexyl aziridine with a dialkyl malonate, whereby to provide a trans-fused 3-alkylcarbonyl-octahydro-indol-2-one; B) decarbonylation at the 3-position, conversion of the ketone of the resulting trans-octahydro-indol-2-one to an optionally protected carboxylic acid group; and C) optionally removing any N-substitution

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if necessary.